

RAW SEQUENCE LISTING ERROR REPORT

The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable form:

Application Serial Number: $\frac{10/803,344}{15400}$ Source: $\frac{1540}{3126/04}$

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.
PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER:

- 1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT, WITH A NOTICE TO COMPLY or,
- 2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT, WITH A NOTICE TO COMPLY

FOR CRF SUBMISSION AND PATENTIN SOFTWARE QUESTIONS, PLEASE CONTACT MARK SPENCER, TELEPHONE: 703-308-4212; FAX: 703-308-4221

Effective 12/13/03: TELEPHONE: 571-272-2510; FAX: 571-273-0221

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE CHECKER VERSION 4.1 PROGRAM, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW FOR ADDRESS:

http://www.uspto.gov/web/offices/pac/checker/chkr41note.htm

Applicants submitting genetic sequence information electronically on diskette or CD-Rom should be aware that there is a possibility that the disk/CD-Rom may have been affected by treatment given to all incoming mail. Please consider using alternate methods of submission for the disk/CD-Rom or replacement disk/CD-Rom.

Any reply including a sequence listing in electronic form should NOT be sent to the 20231 zip code address for the United States Patent and Trademark Office, and instead should be sent via the following to the indicated addresses:

- 1. EFS-Bio (http://www.uspto.gov/ebc/efs/downloads/documents.htm>, EFS Submission User Manual ePAVE)
- 2. U.S. Postal Service: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450
- Hand Carry directly to (EFFECTIVE 12/01/03):
 U.S. Patent and Trademark Office, Box Sequence, Customer Window, Lobby, Room 1B03, Crystal Plaza Two. 2011 South Clark Place, Arlington, VA 22202
- 4. Federal Express, United Parcel Service, or other delivery service to: U.S. Patent and Trademark Office, Box Sequence, Room-4B03-Mailroom, Crystal Plaza Two, 2011 South Clark Place, Arlington, VA 22202

Revised 10/08/03

Raw Sequence Listing Error Summary

ERROR DETECTED	SUGGESTED CORRECTION SERIAL NUMBER: 10/803,344
ATTN: NEW RULES CASES:	PLEASE DISREGARD ENGLISH "ALPHA" HEADERS, WHICH WERE INSERTED BY PTO SOFTWARE
Wrapped Aminos	The number/text at the end of each line "wrapped" down to the next line. This may occur if your file was retrieved in a word processor after creating it. Please adjust your right margin to .3; this will prevent "wrapping."
2Invalid Line Length	The rules require that a line not exceed 72 characters in length. This includes white spaces.
3Misaligned Amino Numbering	The numbering under each 5 th amino acid is misaligned. Do not use tab codes between numbers; use space characters, instead.
4Non-ASCII	The submitted file was not saved in ASCII(DOS) text, as required by the Sequence Rules. Please ensure your subsequent submission is saved in ASCII text.
	Sequence(s) contain n's or Xaa's representing more than one residue. Per Sequence Rules, each n or Xaa can only represent a single residue. Please present the maximum number of each residue having variable length and indicate in the <220>-<223> section that some may be missing.
"bug"	A "bug" in PatentIn version 2.0 has caused the <220>-<223> section to be missing from amino acid sequences(s) Normally, PatentIn would automatically generate this section from the previously coded nucleic acid sequence. Please manually copy the relevant <220>-<223> section to the subsequent amino acid sequence. This applies to the mandatory <220>-<223> sections for Artificial or Unknown sequences.
(OLD RULES)	Sequence(s) missing. If intentional, please insert the following lines for each skipped sequence: (2) INFORMATION FOR SEQ ID NO:X: (insert SEQ ID NO where "X" is shown) (i) SEQUENCE CHARACTERISTICS: (Do not insert any subheadings under this heading) (xi) SEQUENCE DESCRIPTION:SEQ ID NO:X: (insert SEQ ID NO where "X" is shown) This sequence is intentionally skipped
	Please also adjust the "(ii) NUMBER OF SEQUENCES:" response to include the skipped sequences.
8Skipped Sequences (NEW RULES)	Sequence(s) missing. If intentional, please insert the following lines for each skipped sequence. <210> sequence id number <400> sequence id number 000
9Use of n's or Xaa's (NEW RULES)	Use of n's and/or Xaa's have been detected in the Sequence Listing. Per 1.823 of Sequence Rules, use of <220>-<223> is MANDATORY if n's or Xaa's are present. In <220> to <223> section, please explain location of n or Xaa, and which residue n or Xaa represents.
10Invalid <213> Response	Per 1.823 of Sequence Rules, the only valid <213> responses are: Unknown, Artificial Sequence, or scientific name (Genus/species). <220>-<223> section is required when <213> response is Unknown or is Artificial Sequence
11Use of <220>	Sequence(s) missing the <220> "Feature" and associated numeric identifiers and responses. Use of <220> to <223> is MANDATORY if <213> "Organism" response is "Artificial Sequence" or "Unknown." Please explain source of genetic material in <220> to <223> section. (See "Federal Register," 06/01/1998, Vol. 63, No. 104, pp. 29631-32) (Sec. 1.823 of Sequence Rules)
12Patentin 2.0 "bug"	Please do not use "Copy to Disk" function of PatentIn version 2.0. This causes a corrupted file, resulting in missing mandatory numeric identifiers and responses (as indicated on raw sequence listing). Instead, please use "File Manager" or any other manual means to copy file to floppy disk.
13 Misuse of n/Xaa	"n" can only represent a single nucleotide; "Xaa" can only represent a single amino acid



IFWO

RAW SEQUENCE LISTING

DATE: 03/26/2004 TIME: 07:23:05

PATENT APPLICATION: US/10/803,344

Input Set : A:\SYR-HDAC-5005-U.ST25.txt
Output Set: N:\CRF4\03262004\J803344.raw

```
3 <110> APPLICANT: Syrrx, Inc.
    5 <120> TITLE OF INVENTION: HISTONE DEACETYLASE INHIBITORS
   7 <130> FILE REFERENCE: SYR-HDAC-5005-U.
-> 9 <140> CURRENT APPLICATION NUMBER: US/10/803,344
                                                                pr 1-3,5
  10 <141> CURRENT FILING DATE: 2004-03-17
  12 <150> PRIOR APPLICATION NUMBER: US 60/455,437
  13 <151> PRIOR FILING DATE: 2003-03-17
  15 <150> PRIOR APPLICATION NUMBER: US 60/531,203
                                                             Does Not Comply
  16 <151> PRIOR FILING DATE: 2003-12-19
  18 <160> NUMBER OF SEQ ID NOS: 8
                                                        Corrected Diskette Needed
  20 <170> SOFTWARE: PatentIn version 3.2
  22 <210> SEO ID NO: 1
  23 <211> LENGTH: 513
  24 <212> TYPE: PRT (Custom) invalid (2137 response - see item/0 on Ever Surray 25 <213> ORGANISM (Custom) invalid (2137 response - see item/0 on Ever Surray
  28 <220> FEATURE:
-> 29 <221> NAME/KEY: Residues 1-482 of HDAC1 and a 6-histidine tag at the N-terminus
  30 <222> LOCATION: (1)..(513)
  32 <400> SEQUENCE: 1
  34 Met Ser Tyr Tyr His His His His His Asp Tyr Asp Ile Pro Thr
  38 Thr Glu Asn Leu Tyr Phe Gln Gly Ala Met Glu Pro Gly Gly Ser Met
  39 20
                                      25
  42 Ala Gln Thr Gln Gly Thr Arg Arg Lys Val Cys Tyr Tyr Tyr Asp Gly
                                  40
  46 Asp Val Gly Asn Tyr Tyr Gly Gln Gly His Pro Met Lys Pro His
                              55
  50 Arg Ile Arg Met Thr His Asn Leu Leu Leu Asn Tyr Gly Leu Tyr Arg
  51 65
  54 Lys Met Glu Ile Tyr Arg Pro His Lys Ala Asn Ala Glu Glu Met Thr
  58 Lys Tyr His Ser Asp Asp Tyr Ile Lys Phe Leu Arg Ser Ile Arg Pro
         100
                                     105
  62 Asp Asn Met Ser Glu Tyr Ser Lys Gln Met Gln Arg Phe Asn Val Gly
                                 120
  66 Glu Asp Cys Pro Val Phe Asp Gly Leu Phe Glu Phe Cys Gln Leu Ser
                              135
  70 Thr Gly Gly Ser Val Ala Ser Ala Val Lys Leu Asn Lys Gln Gln Thr
```

155

170

150

165

180

74 Asp Ile Ala Val Asn Trp Ala Gly Gly Leu His His Ala Lys Lys Ser

78 Glu Ala Ser Gly Phe Cys Tyr Val Asn Asp Ile Val Leu Ala Ile Leu

185

RAW SEQUENCE LISTING DATE: 03/26/2004 PATENT APPLICATION: US/10/803,344 TIME: 07:23:06

Input Set : A:\SYR-HDAC-5005-U.ST25.txt
Output Set: N:\CRF4\03262004\J803344.raw

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82 Glu Leu Leu Lys Tyr His Gln Arg Val Leu Tyr Ile Asp Ile Asp Ile
               195
    86 His His Gly Asp Gly Val Glu Glu Ala Phe Tyr Thr Thr Asp Arg Val
    90 Met Thr Val Ser Phe His Lys Tyr Gly Glu Tyr Phe Pro Gly Thr Gly
                           230
    94 Asp Leu Arg Asp Ile Gly Ala Gly Lys Gly Lys Tyr Tyr Ala Val Asn
                                          250
                       245
    98 Tyr Pro Leu Arg Asp Gly Ile Asp Asp Glu Ser Tyr Glu Ala Ile Phe
                                      265
    99 260
    102 Lys Pro Val Met Ser Lys Val Met Glu Met Phe Gln Pro Ser Ala Val
                                    280
    103 275
    106 Val Leu Gln Cys Gly Ser Asp Ser Leu Ser Gly Asp Arg Leu Gly Cys
                              295
    110 Phe Asn Leu Thr Ile Lys Gly His Ala Lys Cys Val Glu Phe Val Lys
                            310
                                               315
    114 Ser Phe Asn Leu Pro Met Leu Met Leu Gly Gly Gly Tyr Thr Ile
                                            330
                        325
    118 Arg Asn Val Ala Arg Cys Trp Thr Tyr Glu Thr Ala Val Ala Leu Asp
                    340
    122 Thr Glu Ile Pro Asn Glu Leu Pro Tyr Asn Asp Tyr Phe Glu Tyr Phe
                                    360
               355
    126 Gly Pro Asp Phe Lys Leu His Ile Ser Pro Ser Asn Met Thr Asn Gln
                                                    380
                               375
    130 Asn Thr Asn Glu Tyr Leu Glu Lys Ile Lys Gln Arg Leu Phe Glu Asn
                                                395
                            390
    134 Leu Arg Met Leu Pro His Ala Pro Gly Val Gln Met Gln Ala Ile Pro
                                           410
                       405
    138 Glu Asp Ala Ile Pro Glu Glu Ser Gly Asp Glu Asp Glu Asp Pro
                                                           430
                                        425
                   420
    142 Asp Lys Arg Ile Ser Ile Cys Ser Ser Asp Lys Arg Ile Ala Cys Glu
                                   440
    146 Glu Glu Phe Ser Asp Ser Glu Glu Glu Gly Glu Gly Arg Lys Asn
                                455
            450
    150 Ser Ser Asn Phe Lys Lys Ala Lys Arg Val Lys Thr Glu Asp Glu Lys
                            470
    154 Glu Lys Asp Pro Glu Glu Lys Lys Glu Val Thr Glu Glu Glu Lys Thr
                                            490
                        485
    158 Lys Glu Glu Lys Pro Glu Ala Lys Gly Val Lys Glu Glu Val Lys Leu
                                        505
                   .500
    162 Ala
    166 <210> SEQ ID NO: 2
    167 <211> LENGTH: 1542
    168 <212> TYPE: DNA
                                 sameun
    169 <213> ORGANISM (Custom
    172 <220> FEATURE:
W--> 173 <221> NAME/KEY: DNA sequence encoding residues 1-482 of HDAC1 and a 6-histidine
             tag at the N-terminus
    175 <222> LOCATION: (1)..(1542)
```

RAW SEQUENCE LISTING

PATENT APPLICATION: US/10/803,344

DATE: 03/26/2004 TIME: 07:23:06

Input Set: A:\SYR-HDAC-5005-U.ST25.txt
Output Set: N:\CRF4\03262004\J803344.raw

```
177 <400> SEQUENCE: 2
                                                                                60
    178 atgtcgtact accatcacca tcaccatcac gattacgata tcccaacgac cgaaaacctg
    180 tattttcagg gcgccatgga acccggggga tccatggcgc agacgcaggg cacccggagg
                                                                               120
    182 aaagtotgtt actactacga cggggatgtt ggaaattact attatggaca aggccaccca
                                                                               180
                                                                               240
    184 atgaageete accgaateeg catgaeteat aatttgetge teaactatgg tetetacega
    186 aaaatggaaa tetategeee teacaaagee aatgetgagg agatgaceaa gtaccacage
                                                                               300
    188 gatgactaca ttaaattett gegeteeate egteeagata acatgtegga gtacageaag
                                                                               360
    190 cagatgcaga gattcaacgt tggtgaggac tgtccagtat tcgatggcct gtttgagttc
                                                                               420
    192 tgtcagttgt ctactggtgg ttctgtggca agtgctgtga aacttaataa gcagcagacg
                                                                               480
    194 gacatcgctg tgaattgggc tgggggcctg caccatgcaa agaagtccga ggcatctggc
                                                                               540
                                                                               600
    196 ttctgttacg tcaatgatat cgtcttggcc atcctggaac tgctaaagta tcaccagagg
    198 gtgctgtaca ttgacattga tattcaccat ggtgacggcg tggaagaggc cttctacacc
                                                                               660
    200 acggaccggg tcatgactgt gtcctttcat aagtatggag agtacttccc aggaactggg
                                                                               720
    202 gacctacggg atatcggggc tggcaaaggc aagtattatg ctgttaacta cccgctccga
                                                                               780
    204 gacgggattg atgacgagtc ctatgaggcc attttcaagc cggtcatgtc caaagtaatg
                                                                               840
     206 gagatgttcc agcctagtgc ggtggtctta cagtgtggct cagactccct atctggggat
                                                                               900
     208 cggttaggtt gcttcaatct aactatcaaa ggacacgcca agtgtgtgga atttgtcaag
                                                                               960
     210 agctttaacc tgcctatgct gatgctggga ggcggtggtt acaccattcg taacgttgcc
                                                                              1020
     212 cggtgctgga catatgagac agctgtggcc ctggatacgg agatccctaa tgagcttcca
                                                                              1080
     214 tacaatgact actttgaata etttggacca gatttcaage tecacatcag teettecaat
                                                                              1140
                                                                              1200
     216 atgactaacc agaacacgaa tgagtacctg gagaagatca aacagcgact gtttgagaac
     218 cttagaatgc tgccgcacgc acctggggtc caaatgcagg cgattcctga ggacgccatc
                                                                              1260
                                                                              1320
     220 cctgaggaga gtggcgatga ggacgaagac gaccctgaca agcgcatctc gatctgctcc
     222 tctgacaaac gaattgcctg tgaggaagag ttctccgatt ctgaagagga gggagagggg
                                                                              1380
     224 ggccgcaaga actcttccaa cttcaaaaaa gccaagagag tcaaaacaga ggatgaaaaa
                                                                              1440
     226 gagaaagacc cagaggagaa gaaagaagtc accgaagagg agaaaaccaa ggaggagaag
                                                                              1500
                                                                              1542
     228 ccaqaaqcca aaggggtcaa ggaggaggtc aagttggcct ga
     231 <210> SEQ ID NO: 3
     232 <211> LENGTH: 498
     233 <212> TYPE: PRT/
     234 <213> ORGANISM: Custom
     237 <220> FEATURE:
W--> 238 <221> NAME/KEY: Residues 1-488 of HDAC2 and a 6-histidine tag at the C-terminus
     239 <222> LOCATION: (1)..(498)
     241 <400> SEQUENCE: 3
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                                              10
     244 1
     247 Tyr Tyr Tyr Asp Gly Asp Ile Gly Asn Tyr Tyr Tyr Gly Gln Gly His
     248
                     20
     251 Pro Met Lys Pro His Arg Ile Arg Met Thr His Asn Leu Leu Leu Asn
                                     40
     255 Tyr Gly Leu Tyr Arg Lys Met Glu Ile Tyr Arg Pro His Lys Ala Thr
                                 55
     259 Ala Glu Glu Met Thr Lys Tyr His Ser Asp Glu Tyr Ile Lys Phe Leu
                                                  75
                             70
     260 65
     263 Arg Ser Ile Arg Pro Asp Asn Met Ser Glu Tyr Ser Lys Gln Met Gln
                                              90
                         85
     267 Arg Phe Asn Val Gly Glu Asp Cys Pro Val Phe Asp Gly Leu Phe Glu
                                                              110
                     100
                                          105
     268
```

RAW SEQUENCE LISTING DATE: 03/26/2004 PATENT APPLICATION: US/10/803,344 TIME: 07:23:06

Input Set : A:\SYR-HDAC-5005-U.ST25.txt
Output Set: N:\CRF4\03262004\J803344.raw

271 272	Phe	Cys	Gln 115	Leu	Ser	Thr	Gly	Gly 120	Ser	Val	Ala	Gly	Ala 125	Val	Lys	Leu
275 276		Arg 130		Gln	Thr	Asp	Met 135		Val	Asn	Trp	Ala 140	Gly	Gly	Leu	His
279 280	His		Lys	Lys	Ser	Glu 150		Ser	Gly	Phe	Cys 155	Tyr	Val	Asn	Asp	Ile 160
283 284	Val	Leu	Ala	Ile	Leu 165		Ļeu	Leu	Lys	Tyr 170	His	Gln	Arg	Val	Leu 175	Tyr
287	Ile	Asp	Ile	Asp 180		His	His	Gly	Asp 185		Val	Glu	Glu	Ala 190	Phe	Tyr
288291292	Thr	Thr	Asp		Val	Met	Thr	Val 200		Phe	His	Lys	Tyr 205	Gly	Glu	Tyr
295	Phe	Pro 210		Thr	Gly	Asp	Leu 215		Asp	Ile	Gly	Ala 220		Lys	Gly	Lys
	_		Ala	Val	Asn	Phe 230		Met	Arg	Asp	Gly 235		Asp	Asp	Glu	Ser 240
303	225 Tyr	Gly	Gln	Ile	Phe 245	Lys	Pro	Ile	Ile	Ser 250		Val	Met	Glu	Met 255	
	Gln	Pro	Ser	Ala 260	Val	Val	Leu	Gln	Cys 265		Ala	Asp	Ser	Leu 270		Gly
	Asp	Arg	Leu 275	Gly	Cys	Phe	Asn	Leu 280		Val	Lys	Gly	His 285	Ala	Lys	Cys
	Val		Val	Val	Lys	Thr	Phe 295		Leu	Pro	Leu	Leu 300		Leu	Gly	Gly
		290 Gly	Tyr	Thr	Ile	Arg 310		Val	Ala	Arg	Cys 315	-	Thr	Tyr	Glu	Thr 320
323	305 Ala	Val	Ala	Leu	Asp 325		Glu	Ile	Pro	Asn 330		Leu	Pro	Tyr	Asn 335	Asp
324 327 328	Tyr	Phe	Glu	Tyr 340		Gly	Pro	Asp	Phe 345		Leu	His	Ile	Ser 350		Ser
331	Asn	Met	Thr 355	Asn	Gln	Asn	Thr	Pro 360		Tyr	Met	Glu	Lys 365	Ile	Lys	Gln
		Leu 370	Phe	Glu	Asn	Leu	Arg		Leu	Pro	His	Ala 380		Gly	Val	Gln
	Met	Gln	Ala	Ile	Pro	Glu 390	Asp	Ala	Val	His	Glu 395		Ser	Gly	Asp	Glu 400
343	385 Asp	Gly	Glu	Asp	Pro 405		Lys	Arg	Ile	Ser 410	Ile	Arg	Ala	Ser	Asp 415	Lys
344 347 348		Ile	Ala	Cys 420	Asp	Glu	Glu	Phe	Ser	Asp	Ser	Glu	Asp	Glu 430	Gly	Glu
351	Gly	Gly	Arg 435	Arg	Asn	Val	Ala	Asp	His	Lys	Lys	Gly	Ala 445	Lys	Lys	Ala
	Arg		Glu	Glu	Asp	Lys	Lys 455	Glu		Glu	Asp	Lys 460	Lys		Asp	Val
	Lys		Glu	Asp	Lys	Ser	Lys		Asn	Ser	Gly 475	Glu		Thr	Asp	Thr 480
363		Gly	Thr	Lys		Glu		Leu	Ser	Asn 490	Pro		His	His	His 495	His
364 367		His			485					490					155	

DATE: 03/26/2004

TIME: 07:23:06

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Input Set: A:\SYR-HDAC-5005-U.ST25.txt
                     Output Set: N:\CRF4\03262004\J803344.raw
     371 <210> SEQ ID NO: 4
     372 <211> LENGTH: 1497
     373 <212> TYPE: DNA
     374 <213> ORGANISM: Custom
     377 <220> FEATURE:
W--> 378 <221> NAME/KEY: DNA sequence encoding residues 1-488 of HDAC2 and a 6-histidine
               tag at the C-terminus
W--> 379
     380 <222> LOCATION: (1)..(1497)
     382 <400> SEQUENCE: 4
     383 atgggatcca tggcgtacag tcaaggaggc ggcaaaaaaa aagtetgcta ctactacgac
                                                                                60
     385 ggtgatattg gaaattatta ttatggacag ggtcatccca tgaagcctca tagaatccgc
                                                                               120
     387 atgacccata acttgctgtt aaattatggc ttatacagaa aaatggaaat atataggccc
                                                                               180
     389 cataaagcca ctgccgaaga aatgacaaaa tatcacagtg atgagtatat caaatttcta
                                                                               240
                                                                               300
     391 cggtcaataa gaccagataa catgtctgag tatagtaagc agatgcagag atttaatgtt
     393 ggagaagatt gtccagtgtt tgatggactc tttgagtttt gtcagctctc aactggcggt
                                                                               360
     395 tcagttgctg gagctgtgaa gttaaaccga caacagactg atatggctgt taattgggct
                                                                               420
     397 ggaggattac atcatgctaa gaaatcagaa gcatcaggat tetgttacgt taatgatatt
                                                                                480
     399 gtgcttgcca tccttgaatt actaaagtat catcagagag tcttatatat tgatatagat
                                                                                540
     401 attcatcatg gtgatggtgt tgaagaagct ttttatacaa cagatcgtgt aatgacggta
                                                                                600
     403 tcattccata aatatgggga atactttcct ggcacaggag acttgaggga tattggtgct
                                                                                660
                                                                                720
     405 ggaaaaggca aatactatgc tgtcaatttt ccaatgagag atggtataga tgatgagtca
                                                                                780
     407 tatgggcaga tatttaagcc tattatetea aaggtgatgg agatgtatea acetagtget
     409 gtggtattac agtgtggtgc agactcatta tctggtgata gactgggttg tttcaatcta
                                                                                840
                                                                                900
     411 acagtcaaag gtcatgctaa atgtgtagaa gttgtaaaaa cttttaactt accattactg
                                                                                960
     413 atgcttggag gaggtggcta cacaatccgt aatgttgctc gatgttggac atatgagact
     415 gcagttgccc ttgattgtga gattcccaat gagttgccat ataatgatta ctttgagtat
                                                                              1020
     417 tttggaccag acttcaaact gcatattagt ccttcaaaca tgacaaacca gaacactcca
                                                                              1080
     419 gaatatatgg aaaagataaa acagcgtttg tttgaaaatt tgcgcatgtt acctcatgca
                                                                              1140
     421 cctggtgtcc agatgcaagc tattccagaa gatgctgttc atgaagacag tggagatgaa
                                                                               1200
     423 gatggagaag atccagacaa gagaatttct attcgagcat cagacaagcg gatagcttgt
                                                                               1260
                                                                              1320
     425 gatgaagaat totcagatto tgaggatgaa ggagaaggag gtogaagaaa tgtggotgat
     427 cataagaaag gagcaaagaa agctagaatt gaagaagata agaaagaaac agaggacaaa
                                                                               1380
     429 aaaacagacg ttaaggaaga agataaatcc aaggacaaca gtggtgaaaa aacagatacc
                                                                               1440
     431 aaaggaacca aatcagaaca gctcagcaac cccgggcatc accatcacca tcactaa
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     434 <210> SEQ ID NO: 5
     435 <211> LENGTH: 782
     436 <212> TYPE: PRT
     437 <213> ORGANISM: Custom
     440 <220> FEATURE:
W--> 441 <221> NAME/KEY: Residues 73-845 of HDAC6 and a 6-histidine tag at the C-terminus
     442 <222> LOCATION: (1)..(782)
     444 <400> SEQUENCE: 5
     446 Met Pro Gly Met Asp Leu Asn Leu Glu Ala Glu Ala Leu Ala Gly Thr
                                              10
     447 1
     450 Gly Leu Val Leu Asp Glu Gln Leu Asn Glu Phe His Cys Leu Trp Asp
                                          25
                     20
     451
     454 Asp Ser Phe Pro Glu Gly Pro Glu Arg Leu His Ala Ile Lys Glu Gln
                                      40
     458 Leu Ile Gln Glu Gly Leu Leu Asp Arg Cys Val Ser Phe Gln Ala Arg
```

RAW SEQUENCE LISTING

PATENT APPLICATION: US/10/803,344

The types of errors shown exist throughout the Sequence Listing. Please check subsequent sequences for similar errors.

VERIFICATION SUMMARY

DATE: 03/26/2004 TIME: 07:23:07

PATENT APPLICATION: US/10/803,344

Input Set : A:\SYR-HDAC-5005-U.ST25.txt
Output Set: N:\CRF4\03262004\J803344.raw

L:9 M:270 C: Current Application Number differs, Replaced Current Application Number
L:29 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:1
L:173 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:2
L:174 M:257 W: Feature value mis-spelled or invalid, Describe feature in <223> for SEQ ID#:2
L:238 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:3
L:378 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:4
L:379 M:257 W: Feature value mis-spelled or invalid, Describe feature in <223> for SEQ ID#:4
L:441 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:5
L:649 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:6
L:650 M:257 W: Feature value mis-spelled or invalid, Describe feature in <223> for SEQ ID#:6
L:742 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:7
L:854 M:257 W: Feature value mis-spelled or invalid, <221> Name/Key for SEQ ID#:8
L:855 M:257 W: Feature value mis-spelled or invalid, Describe feature in <223> for SEQ ID#:8